Under the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it contains a valid OMB control number.

Application Data Sheet 37 CFR 1.7			1 76	Attorne	orney Docket Number RBL0		0109-0	01						
				Application Number										
Title of Invention GALACTOMANNANS AND/OR GLUCOMANNANS FOR INCREASING THE BIOAVAILABILITY OF ACTIVE SUBSTANCES														
The application data s bibliographic data arra This document may b document may be prir	nged in a	format specified eted electronically	by the Ur and sub	ited States mitted to the	Patent a	and Tra	demark O	ffice as	outline	ed in 37	CFR 1.	.76.		
Secrecy Ord	er 37	CFR 5.2												
		pplication asso filers only. App												rsuant to
Applicant Inf	orma	tion:						•						
Applicant 1														•
Applicant Autho	rity 💿	nventor OL	egal Re	presentativ	ve unde	r 35	U.S.C. 11	7 ·	OP:	arty of	Interes	t unde	er 35 U.S	S.C. 118
Prefix Given Na			M	liddle Na	ıme			Family Name			· · · · · · · · · · · · · · · · · · ·	Suffix		
Andreas								Hefe	1					
Residence Info	mation	(Select One)	O US	Residen	су (	) No	on US Re	sidenc	у (	) Act	ive US	Milita	ry Servic	e
City Berlingen	0	•	Coun	try Of R	esiden	cei	СН							
Citizenship unde	er 37 C	FR 1.41(b) i	СН								-			
Mailing Address	of App	licant:												
Address 1		Wieslistrasse 3	36								<del></del>			
Address 2									_					
City Berlin	gen					Stat	te/Provir	nce						
Postal Code		CH-8267			Cou	ntryi	СН							
All Inventors Mugenerated within						ation	blocks	may	be			Add		·
Corresponde	ence	Informatio	on:											_
Enter either Cus For further infor					respoi	nden	ce Inforr	natio	n sec	tion t	elow.			
An Address	is bei	ng provided f	or the c	correspo	ndenc	e Info	ormation	of th	is ap	plica	tion.			-
Customer Numb	er	00832												
Email Address jfhoffma@baker				om						Ado	i Email		Remov	e Email
Application	Infor	nation:												
Title of the Inve	Title of the Invention GALACTOMANNANS AND/OR GLUCOMANNANS FOR INCREASING THE BIOAVAILABILITY OF ACTIVE SUBSTANCES							LITY OF						
Attorney Docke	t Numb	er RBL0109-0												
Application Typ	е	Nonprovisi	Nonprovisional											
Subject Matter		Utility	Utility											
Suggested Clas	s (if an	у)				1	Sub Clas	s (if a	any)					
Suggested Tecl	nolog	/ Center (if ar	ıy)			•			•					
Total Number o	Total Number of Drawing Sheets (if any			3			Suggest	ed Fig	jure 1	for Pu	ıblicat	ion (i	if any)	

PTO/SB/14 (08-05)

Approved for use through 07/31/2006. OMB 0651-0032

U.S. Patent and Trademark Office; U.S. DEPARTMENT OF COMMERCE

Under the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it contains a valid OMB control number.

Application Data Sheet 37 CFR 1.76			Attorney D	Docket Number RBL0109-01					
			Application	ion Number			·		
I ITIA AT INVANTIAN I	OMANNANS AND/O NCES	R GLUCOMANNANS FOR INCREASING THE BIOAVAILABILITY OF ACTIVE							
Publication Information	tion:								
Request Early P	ublicati	on (Fee required at	time of Req	uest 37 CFR 1.2	219)				
and certify that t	the inve	I hereby request the tion disclosed in the country, or under the c	the attached	application has	not been and	d will not be t	he subje	ct of	an
Representative Representative informathis information in the A	ation sh	ould be provided fo	r all practitio	ners having a po	ower of attorn	ney in the apation (see 37 (	oplication. CFR 1.32)	Pro	viding
	mer N	lumber or compl	ete the R	epresentative N	lame sectio	n below.			ctions
Please Select One:	(	) Customer Number	O us	Patent Practitions	er O US	S Representati	ive (37 CF	R 11	l .9)
Customer Number	00	0832							
This section allows for tapplication data sheet co. (4), and need not other	constitut	es the specific refere	nce required b						
Prior Application S	tatus	Pending	· · · · · · · · · · · · · · · · · · ·			Ren	nove		
Application Numb	er	Continuity	Туре	Prior Applicat	ion Number	ate (YYYY-MM-DD)			
				10780152		,			
Prior Application S	tatus					Ren	nove		
Application Numb	oer	Continuity	Туре	Prior Application Number Filing Da			ate (YYYY-MM-DD)		
	<u> </u>	a 371 of internationa	ıl	PCT/EP05/01546 2005-02-16			·16		
Additional Domestic the <b>Add</b> button.	Priority	Data may be ge	nerated with	in this form by	selecting		,		
Foreign Priority	y Info	ormation:							
This section allows for not claimed. Providing and 37 CFR 1.55(a).	the appl this info	icant to claim benefit rmation in the applica	of foreign prio	ority and to identify et constitutes the o	any prior fore	ty as required	by 35 U.S	n pric	ority is I 19(b)
A	T	04	i	Boront Cities 5	20to (\\\\\\)	********	nove Driggi		oim od
Application Numb	per	Country i		Parent Filing Date (YYYY-MM-DD)				<u> </u>	aimed
10 2004 008 017.8		DE		2004-03-17	_A! A!		Yes	<u> </u>	No
Additional Foreign P Add button.	riority [	Jata may be gener	ated Within t	nis iorm by sele	cung the				

U.S. Patent and Trademark Office; U.S. DEPARTMENT OF COMMERCE Under the Paperwork Reduction Act of 1995, no persons are required to respond to a collection of information unless it contains a valid OMB control number.

Application Da	sta Shoot 37 CED 1 76	Attorney Docket Number	RBL0109-01		
Application Data Sheet 37 CFR 1.76		Application Number			
Title of Invention GALACTOMANNANS AND/OR GLUCOMANNANS FOR INCREASING THE BIOAVAILABILITY SUBSTANCES					

## **Assignee Information:**

	n in the application data sheet of signment recorded in the Office		rith any requirement of part 3 of Title 37					
Assignee 1								
If the Assignee is an C	Organization check here.	X						
Organization Name	Organization Name Wheli Inter AG							
Mailing Address Info	rmation:							
Address 1	Gartenstrasse 2	Gartenstrasse 2						
Address 2								
City	Zug	State/Province						
Country i CH		Postal Code	CH-6300					
Phone Number		Fax Number						
Email Address								
Additional Assignee D button.	Pata may be generated with	hin this form by selecting the Ac	ld					

## Signature:

A signature of the applicant or representative is required in accordance with 37 CFR 1.33 and 10.18. Please see 37 CFR 1.4(d) for the form of the signature.								
Signature	/John F. Hoffman/			Date (YYYY-MM-DD)	2006-05-04			
First Name	John F. Last Name		Hoffman	Registration Number	26,280			

This collection of information is required by 37 CFR 1.76. The information is required to obtain or retain a benefit by the public which is to file (and by the USPTO to process) an application. Confidentiality is governed by 35 U.S.C. 122 and 37 CFR 1.14. This collection is estimated to take 23 minutes to complete, including gathering, preparing, and submitting the completed application data sheet form to the USPTO. Time will vary depending upon the individual case. Any comments on the amount of time you require to complete this form and/or suggestions for reducing this burden, should be sent to the Chief Information Officer, U.S. Patent and Trademark Office, U.S. Department of Commerce, P.O. Box 1450, Alexandria, VA 22313-1450. DO NOT SEND FEES OR COMPLETED FORMS TO THIS ADDRESS. SEND TO: Commissioner for Patents, P.O. Box 1450, Alexandria, VA 22313-1450.

## **Privacy Act Statement**

The Privacy Act of 1974 (P.L. 93-579) requires that you be given certain information in connection with your submission of the attached form related to a patent application or patent. Accordingly, pursuant to the requirements of the Act, please be advised that: (1) the general authority for the collection of this information is 35 U.S.C. 2(b)(2); (2) furnishing of the information solicited is voluntary; and (3) the principal purpose for which the information is used by the U.S. Patent and Trademark Office is to process and/or examine your submission related to a patent application or patent. If you do not furnish the requested information, the U.S. Patent and Trademark Office may not be able to process and/or examine your submission, which may result in termination of proceedings or abandonment of the application or expiration of the patent.

The information provided by you in this form will be subject to the following routine uses:

- 1. The information on this form will be treated confidentially to the extent allowed under the Freedom of Information Act (5 U.S.C. 552) and the Privacy Act (5 U.S.C. 552a). Records from this system of records may be disclosed to the Department of Justice to determine whether the Freedom of Information Act requires disclosure of these records.
- A record from this system of records may be disclosed, as a routine use, in the course of presenting evidence to a court, magistrate, or administrative tribunal, including disclosures to opposing counsel in the course of settlement negotiations.
- 3. A record in this system of records may be disclosed, as a routine use, to a Member of Congress submitting a request involving an individual, to whom the record pertains, when the individual has requested assistance from the Member with respect to the subject matter of the record.
- 4. A record in this system of records may be disclosed, as a routine use, to a contractor of the Agency having need for the information in order to perform a contract. Recipients of information shall be required to comply with the requirements of the Privacy Act of 1974, as amended, pursuant to 5 U.S.C. 552a(m).
- 5. A record related to an International Application filed under the Patent Cooperation Treaty in this system of records may be disclosed, as a routine use, to the International Bureau of the World Intellectual Property Organization, pursuant to the Patent Cooperation Treaty.
- 6. A record in this system of records may be disclosed, as a routine use, to another federal agency for purposes of National Security review (35 U.S.C. 181) and for review pursuant to the Atomic Energy Act (42 U.S.C. 218(c)).
- 7. A record from this system of records may be disclosed, as a routine use, to the Administrator, General Services, or his/her designee, during an inspection of records conducted by GSA as part of that agency's responsibility to recommend improvements in records management practices and programs, under authority of 44 U.S.C. 2904 and 2906. Such disclosure shall be made in accordance with the GSA regulations governing inspection of records for this purpose, and any other relevant (i.e., GSA or Commerce) directive. Such disclosure shall not be used to make determinations about individuals.
- A record from this system of records may be disclosed, as a routine use, to the public after either publication of the application pursuant to 35 U.S.C. 122(b) or issuance of a patent pursuant to 35 U.S.C. 151. Further, a record may be disclosed, subject to the limitations of 37 CFR 1.14, as a routine use, to the public if the record was filed in an application which became abandoned or in which the proceedings were terminated and which application is referenced by either a published application, an application open to public inspections or an issued patent.
- 9. A record from this system of records may be disclosed, as a routine use, to a Federal, State, or local law enforcement agency, if the USPTO becomes aware of a violation or potential violation of law or regulation.